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Shaping RNA Structures with Metal Ions and Metal Ion Complexes

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[§]Werner Prize Winner 2009

Abstract: The research in our laboratory focuses on the role of metal ions and their complexes in structure formation and folding of nucleic acids. Large catalytic RNAs, like group II introns and some riboswitches, as well as shorter RNAs and DNAs containing modified nucleotides for the assembly of nanodevices are examined. Abundant metal ions like Mg^{2+} or natural metabolites like coenzyme B_{12} are in the center of interest, but also other metal ions, complexes thereof and B_{12} derivatives are applied with the aim to understand the largely unknown and manifold non-covalent interactions with nucleic acids. We apply a multitude of techniques, including potentiometric pH titrations, NMR spectroscopy, X-ray crystallography, gel electrophoresis and single molecule FRET experiments. Here we briefly summarize each of our research topics emphasizing the interaction of coenzyme B_{12} and its derivatives with the *btuB* riboswitch of *E. coli*. This highly conserved sequence, found in the 5'-untranslated region (5'-UTR) of the *btuB* mRNA, is involved in the regulation of the *btuB* protein expression. After a summary on the historical discovery of such riboswitches and their mechanism of action, we shortly focus on our own contributions to understand the structural equilibrium, high affinity and selectivity of the interaction between this specific RNA sequence and the largest and most complex cellular metabolite, coenzyme B_{12} .

Keywords: Coenzyme B_{12} · Metal ions · Ribozymes · Riboswitches · RNA/DNA



Roland K. O. Sigel is Associate Professor (2009) of Inorganic Chemistry at the University of Zurich. He studied chemistry at the University of Basel, received his PhD *summa cum laude*

(1999) from the University of Dortmund, and thereafter spent nearly three years at Columbia University, New York. He has received several fellowships, was endowed with a SNF-Förderungsforschung (2003–2008), invited to plenary lectures at several international conferences, and in 2009 he received a Lectureship of the Hungarian Chemical Society. In 2008 he has been awarded the *EuroBIC Medal* and in 2009 the *Werner Prize* of the Swiss Chemical Society. His research focuses on

the characterization of metal ion-nucleic acid interactions, and he is a co-editor of the series *Metal Ions in Life Sciences*.

1. Introduction

Since the discovery of catalytically active RNA sequences, named *ribozymes* from here on, by the Cech and Altman laboratories in the early 1980s,^[1,2] the knowledge on functional RNAs, being either synthetic or natural, has steadily increased. Today it is known that RNA is involved in a multitude of cellular processes, sometimes also performing catalytic functions. RNA molecules can thereby act either independently of any protein cofactor or as RNA-protein complexes, e.g. the ribosome (protein biosynthesis) or the spliceosome (mRNA splicing in eukaryotic cells).^[3]

Independent of the nucleic acids' size and function, metal ions are the most crucial and abundant cofactors, determining folding, global, and local structures, as well as taking directly part in catalysis of ribozymes.^[4] Mg^{2+} and K^+ are usually considered the naturally involved metal ions with nucleic acids in the cell.^[5] However, also other metal ions are available and might take part in folding and catalysis.^[5–8] In all cases it is unknown so far how metal ions exactly influence the structure and activity on the atomic level. To answer this fundamental question is the driving force of our research.

2. Metal ions in the Catalytic Center of Group II Intron Ribozymes

Group II introns are natural self-splicing catalytic RNAs and are among the largest ribozymes known.^[9] They occur in some bacteria, but mainly in plants, fungi, and lower eukaryotes. These macromolecules are closely related to the eukaryotic splicing machinery and are considered the ancestors of up to about 40% of the human genome.^[10] Group II introns catalyze a diverse set of reactions like splicing but also reinsertion into DNA, being mobile genetic elements. The nature of the metal ion present in solution has thereby a strong influence on the catalytic activity of these ribozymes.^[8,11–13] Recently, a first X-ray structure of a self-splicing group II intron has been solved, although unfortunately at too low resolution to learn much about the metal ions present.^[14]

We aim to determine the specific metal ion binding sites within the active structure and characterize their thermodynamic properties. By means of NMR, we determine the solution structures of domains and subdomains which are part of the catalytic core and subsequently characterize their metal ion binding properties by various spectroscopic techniques (Fig. 1).^[15–22] One of the biggest challenges is the determination of the metal ion affinities because usually several metal ions simultaneously bind to one RNA molecule with similar affinities. We have thus developed

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an iterative approach to calculate the intrinsic binding constants of different metal ions to one ligand, *e.g.* a RNA hairpin, based on NMR titration experiments.^[17] Further NMR experiments are conducted to identify the specific liganding atoms of the RNA;^[18] The main problem hereby lies in the fact that each of the four nucleotide building blocks offers a multitude of potential coordination sites, *e.g.* the phosphate oxygens or the N(7) of the purines. At the same time inner-sphere, *i.e.* direct coordination, or outer-sphere binding, mediated through a water molecule, can take place, further complicating the situation. Nevertheless, a surprising selectivity and specificity of RNAs for given metal ions is reached,^[8] which we aim to reveal on the coordination chemical level (for a recent review, see ref. [21]). Only in this way will we finally be able to understand the ribozyme mechanism and modify these molecules for further needs.

3. From M^{n+} Binding to Nucleotides to Nucleic Acids

Metal ion coordination to mononucleotides is relatively well understood.^[24–26] In the past years, we have extended these studies towards phosphorothioate derivatives,^[27,28] dinucleotides,^[29–31] as well as the binding of two metal ions to one nucleobase.^[32] Surprisingly, in the latter case, the binding of the second metal ion is only slightly hampered by the presence of the first one.^[32] We have now used this accumulated wealth of information to create a library of stability increments for the individual liganding atoms towards Mg^{2+} , Ca^{2+} , Mn^{2+} , Cu^{2+} , Zn^{2+} , Cd^{2+} or Pb^{2+} .^[33] These increments can be applied to estimate the intrinsic affinities of nucleic acid-binding pockets for the mentioned metal ions and thus to provide first insights into the thermodynamics of individual binding pockets within large nucleic acid structures.

To assist the above-mentioned approach of adding up stability increments and to allow for a rigorous examination of all known metal ion binding pockets, we have built up a freely available database on *Metal Ions in Nucleic Acids* (MINAS) (Fig. 2).^[34] This database can be used to evaluate, visualize, and characterize the metal ion binding sites of all structures deposited in the PDB and NDB.

4. Metal Ion-assisted Folding of Large RNAs

Metal ions induce and direct the folding of RNAs.^[35,36] In most cases, the presence of divalent metal ions leads to a rapid collapse into different kinetic traps and to

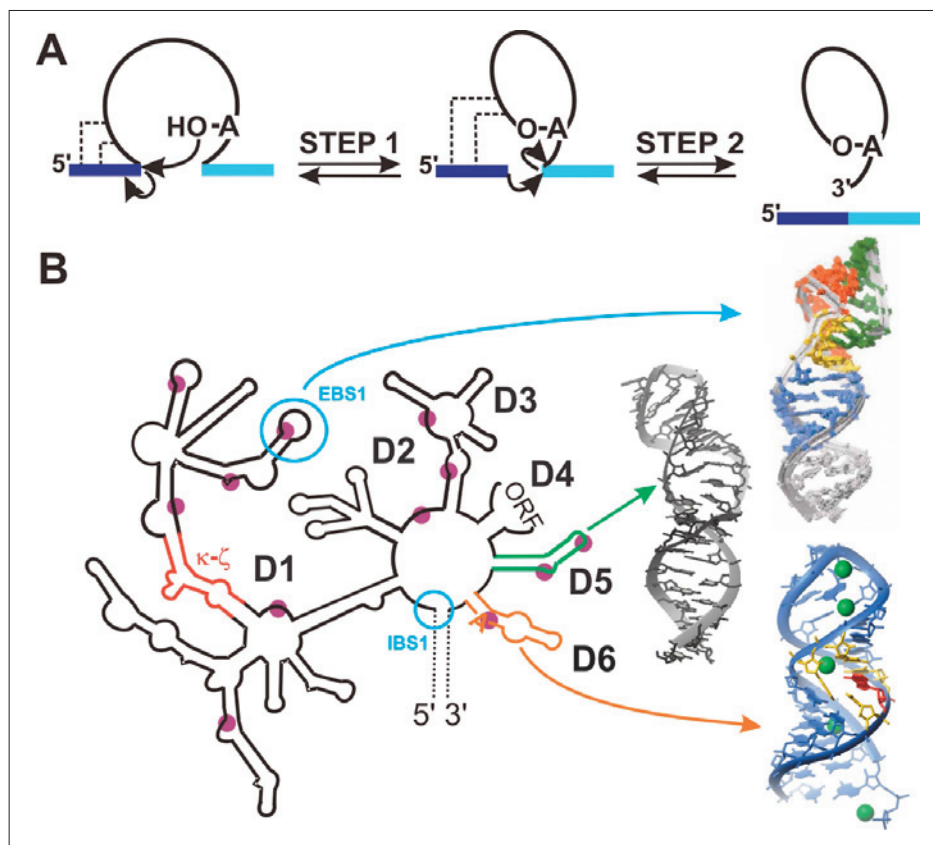


Fig. 1. Group II intron ribozymes. (A) Schematic splicing reaction, *i.e.* the removal of the intronic sequence and joining of the two flanking exons. Both steps are reversible and performed without the aid of proteins. (B) Schematic secondary structure of the yeast mitochondrial group II intron Sc.ai5γ. Regions that are part of the catalytic core and in the focus of our research are colored and metal ion binding sites indicated as magenta spheres. NMR solution structures of the intron-exon interaction, domains 5 and 6, as solved in our lab, are shown to the right, based on the PDB files 2K64,^[20] 1R2P,^[15] and 2AHT^[16] and prepared with MOLMOL.^[23] Domain 5 is the most conserved region of the intron and domain 6 contains the nucleophile in the first step of splicing.

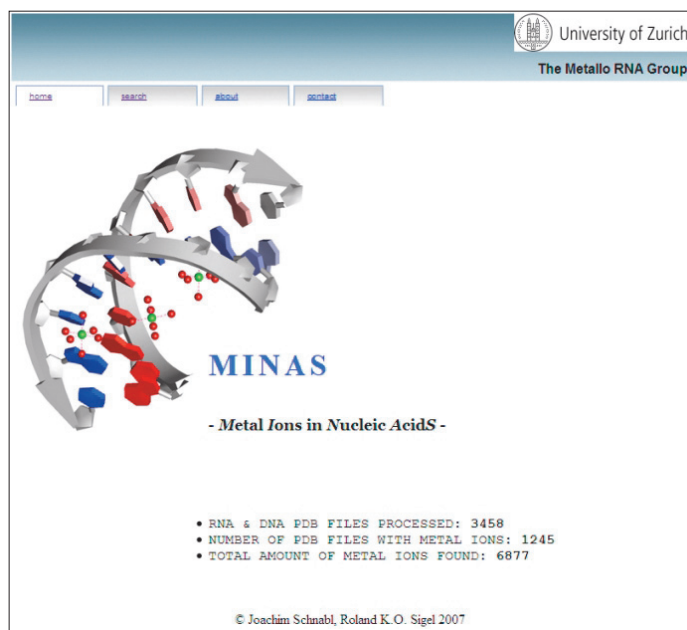


Fig. 2. Database of Metal Ions in Nucleic Acids (MINAS; www.minas.uzh.ch). This database contains all metal ion-nucleic acid structures of the PDB and NDB and allows a detailed analysis of the metal ion coordination spheres.

a subsequent slow three-dimensional rearrangement of the (sub)domains into the active structure.^[37,38] Group II introns behave differently: A first slow folding step is followed by a series of fast consecutive

steps obeying a straight pathway devoid of kinetic traps.^[39–41] We have recently characterized the folding pathway of one of these large RNAs by single molecule Fluorescence Resonance Energy Transfer

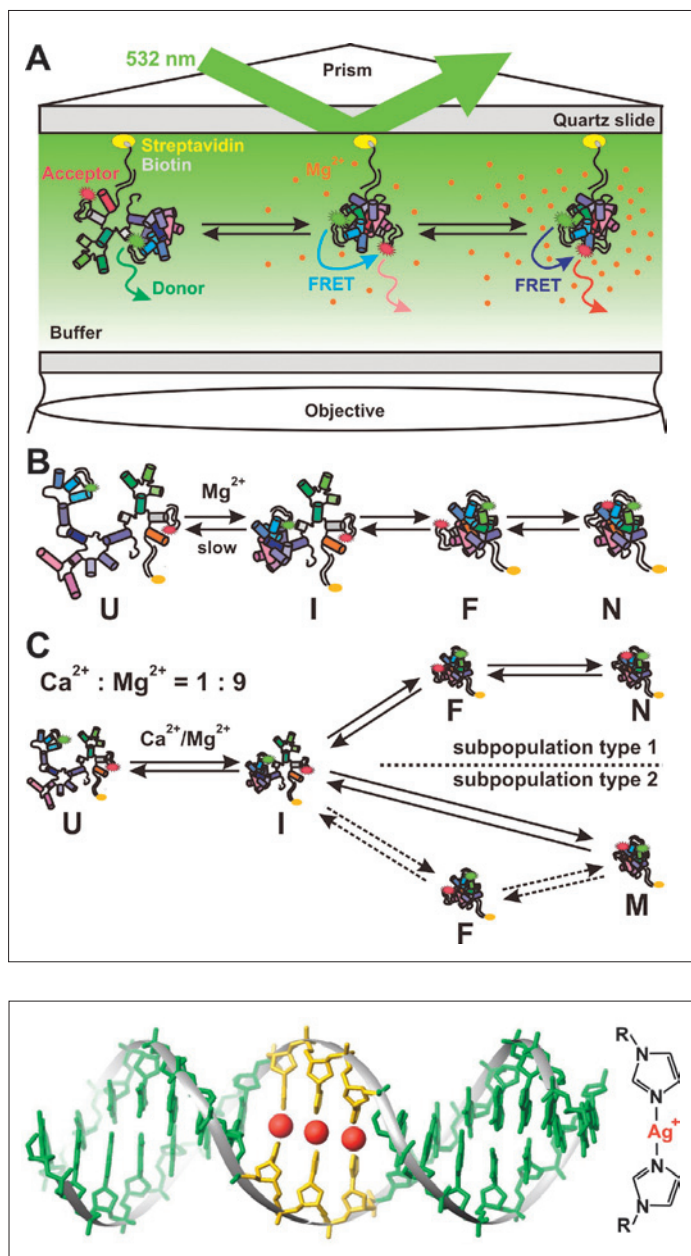


Fig. 3. Single molecule FRET studies of group II intron folding. (A) Schematic setup of smFRET experiments: The RNA molecules are attached via a biotin/streptavidin interaction to a quartz slide, the donor fluorophore excited by a laser, and the fluorescence measured by a CCD camera from below. (B) Folding pathway of a *Sc.ai5 γ* derived ribozyme construct in the presence of Mg^{2+} only being devoid of kinetic traps (U, unfolded; I, intermediate, F, folded; N, native state). (C) Folding pathway in the presence of Ca^{2+} . The molecules fold into two distinct subpopulations that do not interchange (U, unfolded; I, intermediate, F, folded; M, misfolded; N, native state).

Fig. 4. First NMR solution structure of a metal-modified DNA duplex containing three imidazole- Ag^+ -imidazole base pairs in its center, as indicated on the right. The Ag^+ ions are shown by red spheres. The Fig. has been prepared based on the structure 2KE8^[46] using MOLMOL.^[23]

(smFRET) spectroscopy^[42] and investigated the effect of Ca^{2+} on this folding pathway (Fig. 3);^[43] Already small amounts of Ca^{2+} affect the global structure of this ~700 nucleotide long RNA. Unprecedentedly, two distinct subpopulations were observed that do not interchange with each other. It was highly surprising to see the single molecules fold into two distinctly different three-dimensional structures occurring in parallel under the same conditions.^[43]

5. From Nature to Nano-devices

Metal ions usually bind to the major and minor groove edges of nucleotides within

duplex or complex nucleic acids as they occur manifold in nature. Our interests lie in the investigation of structures containing metal ions in the center of the nucleic acid duplex. For example, Hg^{2+} ions can be 'trapped' in the helix center between two opposite thymine or uracil nucleobases, which become deprotonated upon coordinating the metal ion at N(3).^[44,45] Recently we succeeded in determining the first three-dimensional structure of a metallated DNA duplex containing artificial imidazole nucleotides with three linearly coordinated Ag^+ ions aligned in the middle (Fig. 4).^[46] Based on solution structures and investigations on the metal ion-initiated structural changes as performed

in our laboratory, the aim is to construct nanodevices with particular electric, magnetic and sensing properties.^[47,48]

6. Sensing Metal Ion Metabolites by RNA

Riboswitches are recently discovered highly conserved sequences in the untranslated regions (UTR) of mostly bacterial mRNAs. Upon binding to certain metabolites with high specificity and selectivity these RNAs change their three-dimensional structure thus regulating gene expression and metabolite concentration on the translational or transcriptional level.^[49] Our research focuses on the Mg^{2+} - and coenzyme B_{12} -sensitive riboswitches^[50,51] whereby the latter is explained in more detail in the following Section.

7. Riboswitches

Riboswitches are conserved sequences within mRNA guiding very elegantly protein-independent gene-regulation.^[49,51–54] Out of the more than 20 riboswitch classes discovered so far, only the TPP-binding riboswitch is known to be present also in archaea, fungi and plants,^[55–58] whereas all others are restricted to bacteria where they are believed to regulate up to 2% of the genes.^[59] Due to their overwhelming occurrence in bacteria, riboswitches are of increasing interest as new targets for the design of antimicrobial drugs.^[60]

Riboswitches act as natural aptamers that bind to small metabolites with very high affinity. Such metabolites are amino acids, purine bases, redox cofactors, metal ions, second messengers and others. It is highly intriguing how RNA composed of only four building blocks is able to recognize and bind such a variety of ligands different in size, charge and functional groups. As it turns out, hydrogen-bonding, electrostatic and van der Waals interactions as well as Watson-Crick base pairing and stacking interactions are used concomitantly to ensure specificity and affinity.^[56,61–63]

Riboswitches generally consist of a highly conserved aptamer domain, a region of 35 to 260 nucleotides, which represents the metabolite binding part, and a downstream expression platform. Metabolite binding causes considerable conformational changes in secondary and tertiary structure and subsequent reorganization of the expression platform, the latter being finally responsible for gene regulation. The downstream genes all have a close physiological relationship with the bound metabolite, *i.e.* they encode for transport proteins or proteins involved in the biosyn-

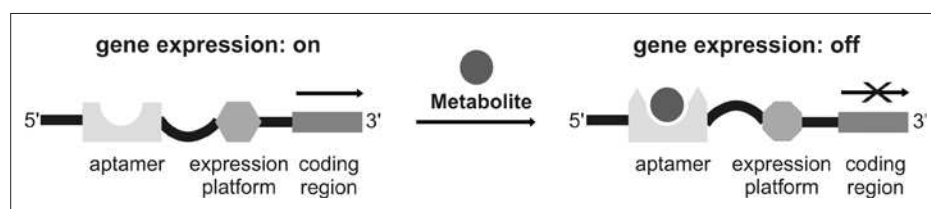


Fig. 5. General mechanism of a riboswitch. Binding of the metabolite to its aptamer region of the riboswitch leads to a structural rearrangement of this domain as well as of the expression platform and subsequently to an altered gene expression. Such metabolites are amino acids (lysine and glycine), purine bases (adenine, guanine and prequenosine-1), cofactors (AdoCbl, flavin mononucleotide, S-adenosyl methionine, tetrahydrofolate, thiamine pyrophosphate, molybdenum and tungsten cofactors), metal ions (Mg^{2+}), small nucleotide-units acting as second messenger (c-di-GMP) or others (glucosamine-6-phosphate, S-adenosyl homocysteine).^[49]

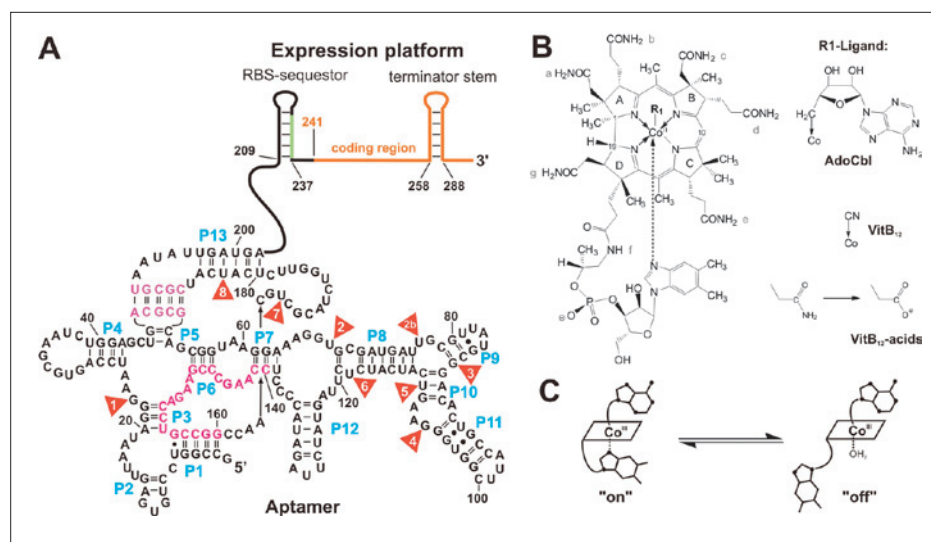


Fig. 6. The *btuB* riboswitch and its metabolite coenzyme B_{12} . (A) Secondary structure of the *btuB* riboswitch from *E. coli* in the presence of AdoCbl. The aptamer region of the riboswitch covers nucleotide 1 to 202 and includes a pseudoknot (magenta) and the B_{12} -box (pink).^[71] The expression platform is situated between nucleotides 209 and 288 and contains a translational regulatory element (the ribosomal binding site(RBS)-sequester hairpin with the Shine-Dalgarno sequence in light green)^[72] and a transcriptional regulatory element (termination stem) being part of the coding sequence (orange).^[73] (B) General formula of B_{12} molecules with an axial dimethyl benzimidazole ligand as present in AdoCbl, Vit B_{12} , and their acid derivatives. (C) Intramolecular equilibrium between the 'base-on' and 'base-off' forms of the B_{12} derivatives. Under physiological conditions, AdoCbl is found exclusively in the 'base-on' form, whereas in the case of adenosyl factor A the 'base-off' form prevails.

thesis of the metabolite (Fig. 5). Regulation can thereby occur based on structural changes, self-cleavage^[64] or (alternative) splicing.^[49,57] Genetic control is mostly negative (*i.e.* switching off gene expression) but there are also some examples of a positive feedback such as the adenine responding *ydhl* riboswitch.^[65] In addition, cooperativity^[66] or multiple recognition units within so-called tandem riboswitches are known.^[67,68]

7.1 Discovery and Structures of B_{12} Responsive RNAs

B_{12} sensing riboswitches are widespread throughout the bacterial kingdom and are found in nearly all taxonomic groups. They control gene products involved in cobalamin synthesis and trans-

port, cobalt transport, B_{12} -independent aerobic and anaerobic ribonucleotide reductase synthesis, as well as glutamate and succinate fermentation.^[69]

Kadner and coworkers recorded the first structural influence of vitamin B_{12} (Vit B_{12}) and adenosyl cobalamin (coenzyme B_{12} , AdoCbl) on a naturally occurring RNA in 1991 with the *btuB* leading sequence of *E. coli* (Fig. 6A).^[70] Subsequent investigations with the *btuB* gene and *cob* operon of *S. typhimurium* revealed a highly conserved 25-nt long sequence essential for gene regulation and high-level expression called the B_{12} -box.^[73,74] By reverse transcription three highly structured regions were identified within this RNA:^[72] i) the ribosome binding site at Thy256, ii) Ade202 marking the start of

the *btuB* aptamer region, and iii) Cyt161 at the beginning of the B_{12} box. Whereas the latter structure is independent of AdoCbl, the Ade202 region showed a clear structural dependence on AdoCbl concentration. In 2003, a sequence alignment of approximately 200 B_{12} -aptamers revealed a high degree of sequence conservation and a first common secondary structure could be proposed.^[75] This consensus sequence included six hairpins arranged around a central wheel as well as an antiterminator or/and an antisequestor hairpin belonging to the expression platform.

Further investigations revealed two types of B_{12} aptamers: The first includes the *btuB* riboswitches and the longer *cob* operon leading sequence with at least ten base-paired regions and around 57 conserved nucleotides. The second class was found in *B. subtilis*, upstream of the *yvrC* gene lacking the large part around helices P8 to P12.^[71]

7.2 Gene Regulation by the B_{12} Sensing Riboswitches

Depending whether an organism can synthesize or has to take up coenzyme B_{12} , B_{12} -dependent riboswitches control genes involved in the biosynthesis of B_{12} (*e.g.* *S. typhimurium*) or uptake and transport of the same (*e.g.* *E. coli*). For example, the *cob* operon of *S. typhimurium* encodes 20 proteins of the cobalamin biosynthetic pathway and is controlled by the *cob* riboswitch.^[76,77] Instead, the *btuB* riboswitch regulates a single gene encoding for an outer membrane transporter responsible for the active transport of cobalamins into the periplasmic space.^[78–80] Both riboswitches are negatively controlled on both the transcriptional and the translational level and the structural rearrangements seem to correlate well.^[70,81] In the *btuB* riboswitch, the first 240 nucleotides are responsible for the translational control of gene expression by sequestration of the Shine-Dalgarno sequence, the binding site of the 16S rRNA unit of the ribosome (Fig. 6A). Instead, the first 100 nucleotides of the coding sequence are compulsory for a Rho-independent transcriptional regulation (Fig. 6).^[73] A possible mechanism of gene regulation is the formation of a pseudo-knot upon binding to AdoCbl and the resulting reorganization of the expression platform. However in-line probing experiments demonstrated also a substantial structural rearrangement of the aptamer region upon binding to AdoCbl and some of its derivatives (Fig. 7).^[53]

7.3 Specific Recognition of AdoCbl by the *btuB* Riboswitch

The organometallic complex coenzyme B_{12} is the largest and most complex of all coenzymes (Fig. 6B). AdoCbl and its

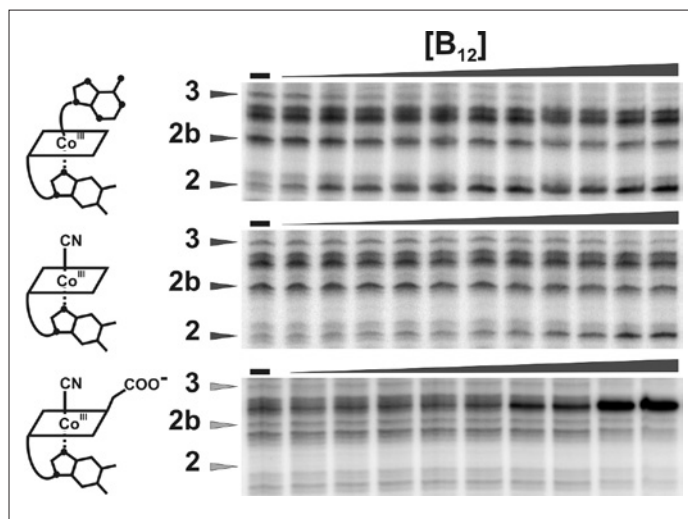


Fig. 7. In-line probing experiments with the *btuB* riboswitch in the presence of different B_{12} derivatives. Addition of AdoCbl (top) leads to the decrease of the intensity of bands 3 and 2b while cleavage increases at site 2. Vit B_{12} (middle) shows the same pattern but only at higher concentrations. *c*-acid Vit B_{12} (bottom) binds, but prevents the riboswitch from a correct rearrangement, as is obvious from the different cleavage pattern of the RNA. B_{12} concentrations range from 30 nM to 0.1 mM (AdoCbl), 3.3 μ M to 10 mM (Vit B_{12}), and 1 μ M to 0.75 mM (*c*-acid Vit B_{12}).

derivatives are considered the evolutionary most ancient protein cofactors^[82–84] and might have already played a role in the so-called RNA world.^[85] This ancient relationship between a nucleic acid and this fascinating metal ion complex triggered our interest already some time ago.

The first example of a RNA that binds specifically to a B_{12} derivative was discovered in 1994 by Szostak and coworkers in *in vitro* selection experiments.^[86] This synthetic aptamer of 35 nucleotides in length selectively binds Vit B_{12} ($K_D = 90$ nM) over other B_{12} derivatives such as dicyanocobinamide (20 mM) or the larger AdoCbl, which does not bind. The X-ray crystal structure^[87] revealed that a large hydrophobic pocket is situated at the interface between a small duplex and a unique triple helix structure. Selective Vit B_{12} binding is achieved by a dense network of different interactions including hydrophobic and electrostatic interactions, hydrogen-bonding as well as the insertion of the cyanide ligand of Vit B_{12} into the base-triplet of the triple helix, the latter explaining the selectivity for cyanide-carrying B_{12} molecules.

As numerous B_{12} derivatives occur in nature, a different selection scheme is expected for natural B_{12} aptamers. Indeed, the two best studied B_{12} riboswitches, the *btuB* riboswitch and the leading sequence of the *cob* operon, show a high selectivity for AdoCbl (the most abundant form of B_{12} , $K_D = 89 \pm 6$ nM with *btuB*) over Vit B_{12} ($K_D = 314 \pm 141$ μ M).^[50,53,81] Our investigations concentrate currently on the

structural basis for the interaction between AdoCbl and the 202 nt long aptamer region of the *btuB* riboswitch of *E. coli*.^[51] In contrast to Vit B_{12} , AdoCbl is able to release the lower dimethylbenzimidazol ligand in an intramolecular equilibrium, *i.e.* leading to a ‘base-off’ form (Fig. 6C). From protein interactions it is known that either the ‘base-on’ or the ‘base-off’ form can be bound.

In order to elucidate the binding mode to the riboswitch, we applied various structurally different B_{12} derivatives, *i.e.* missing either the upper (Vit B_{12}), the lower (adenosyl cobinamide) or both large apical ligands (dicyanocobinamide), or displaying the ‘base-off’-form (adenosyl factor A).^[51] The conformational changes of the RNA upon interaction with the different B_{12} derivatives was visualized by in-line probing experiments, *i.e.* structure-dependent self-degradation of RNA (Fig. 7). The conformational switch of the RNA was followed by the appearance or disappearance of nine distinct cleavage bands upon binding to AdoCbl (Fig. 7).^[51,53] The nine cleavage sites are spread over the whole sequence, implying that binding to AdoCbl leads to an overall structural rearrangement of the RNA.

In contrast to earlier studies,^[53,71,72] our experiments showed the adenosyl-moiety on the upper side of the corrin ring not to be compulsory for the switching of the RNA.^[51] If present in high concentrations, also Vit B_{12} induces a correct switch of the *btuB* RNA (Fig. 7). Similarly, the pres-

ence and conformation of the lower part of the B_{12} derivative is of minor importance. Adenosyl cobinamide as well as the sterically more demanding adenosyl factor A (AdoFactA) allow essentially the same rearrangement of the riboswitch as AdoCbl. The unexpected high affinity for AdoFactA ($K_D = 290 \pm 32$ nM) further suggests that the ‘base-off’ nature of this ligand does not disturb the interaction with the RNA. UV studies showed further that the lower base is not replaced by a RNA nucleotide at the cobalt coordination site.^[51] As a direct consequence of the indicated results, one can conclude that the presence of the two axial ligands increase the B_{12} -RNA interaction each by a factor of almost 1000.^[51]

In contrast, the corrin ring with its side chains is directly responsible for the structural switch of the *btuB* RNA. We investigated the role of single amide side chains on riboswitch restructuring by applying B_{12} derivatives such as the *b*-, *c*-, *d*- and *e*-acid derivatives of Vit B_{12} (Fig. 6B) in in-line probing experiments. Application of these derivatives, all having one single amide group replaced by a carboxylic acid, provides a first insight in the importance of the correct H-bonding pattern and electrostatic environment on *btuB* RNA switching.^[50] Interestingly, most B_{12} -acid derivatives only lead to a partial rearrangement, whereas binding of the *c*-acid B_{12} is followed by a completely altered structural change. These findings corroborate the hypothesis that the amide side chains of the corrin ring are of uttermost importance for *btuB* RNA switching.

8. Conclusions and Outlook

The specific coordination of metal ions and metal ion complexes to nucleic acids is highly complex and goes in most instances far beyond a simple coulombic interaction. Relatively fast ligand exchange rates, rather low affinities, and the mostly spectroscopically silent metal ions involved, make the characterization of this interplay highly challenging.^[5,8] The combination of inner-sphere metal ion coordination with an extensive hydrogen bonding network *via* coordinated water molecules allows on the one hand a rather tight binding in preformed pockets, but on the other hand also a high degree of flexibility and dynamics. The dynamics of metal ion binding seem to play a crucial role for ribozyme activity, *e.g.* single group II intron molecules become more dynamic in their domain movements with increasing Mg^{2+} concentration.^[42]

Our efforts aim to elucidate the basic principles of metal ion binding and to characterize the effect thereof on global structure, dynamics, and catalytic mechanism.

The importance of the knowledge of, e.g. the intrinsic binding affinities is exemplified by the recent discovery that the catalytic rate of the hammerhead ribozyme is directly dependent on the phosphate affinity of the divalent metal ion present.^[8] Such relatively 'simple' correlations might turn out to be also true for other small ribozymes. In the case of the large ribozymes, more complicated interdependencies between metal ions and the RNA are to be expected because of the above-mentioned reasons.

To conclude, a highly interdisciplinary approach will be needed to understand the diverse world of ribozymes and ribo-switch structures and functions depending on metal ions and/or metal ion complexes. At the same time we may expect surprising and unprecedented turns and twists in this fascinating field of *Biological Inorganic Chemistry*.

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